CENTER FOR DRUG EVALUATION AND RESEARCH

APPLICATION NUMBER: 40333

BIOEQUIVALENCY REVIEW(S)

OFFICE OF GENERIC DRUGS

DIVISION OF BIOEQUIVALENCE

ANDA # 40-333 SPONSOR: Gensia Sicor Pharmaceutical, Inc.

DRUG & DOSAGE FORM:	Fluoroura	cil Inject:	ion	,			
STRENGTH: 50 mg/mL,	500 mg in	10 mL via	1				
TYPE OF STUDY: SD	SDF	MULT	OTHER	Waiver Request			
STUDY SITE: NA	CLINICAL:	NA	ANALYT	ICAL: NA			
STUDY SUMMARY: The waiver of <i>in vivo</i> bioequivalence study is granted per 21 CFR § 320.22(b)(1) of Bioavailability/Bioequivalence Regulations.							
PRIMARY REVIEWER; Chandra S. Chaurasia, Ph.D. BRANCH: I DATE: 11/2/98							
TEAM LEADER: Yih Cha	_			BRANCH: I DATE: $11/2/98$			
DIRECTOR, DIVISION O				Conner, Pharm.D. DATE: ///2/98			
DIRECTOR, OFFICE OF INITIAL:				DATE:			

BIOEQUIVALENCY COMMENTS

ANDA: #40-333 APPLICANT: Gensia Sicor Pharmaceutical, Inc

DRUG PRODUCT: Fluorouracil Injection, USP 50 mg/mL; 10 mL Vial

The Division of Bioequivalence has completed its review of your application and has no further questions at this time.

Please note that the bioequivalency comments provided in this communication are preliminary. These comments are subject to revision after review of the entire application, upon consideration of the chemistry, manufacturing and controls, microbiology, labeling, or other scientific or regulatory issues. Please be advised that these reviews may result in the need for additional bioequivalency information and/or studies, or may result in a conclusion that the proposed formulation is not approvable.

Sincerely yours,

Dale P. Conner, Pharm. D.

Director, Division of Bioequivalence

Office of Generic Drugs

Center for Drug Evaluation and Research

CC: ANDA #40-333 ANDA DUPLICATE

DIVISION FILE

HFD-650/Bio Secretary-Bio Drug File

HFD-652/C. Chaurasia

Endorsements:

HFD-652/C. Chaurasia (* 11/2/98

HFD-652/YC Huang 4 1/2/78 HFD-650/D. Conner 8 1/2/98

BIOEQUIVALENCY - ACCEPTABLE

WAIVER (WAI)

Strength: 50mg/mL

Outcome: AC

Outcome Decisions: Acceptable

AC - Acceptable

WINBIO COMMENTS: The waiver is granted

Fluorouracil Injection, USP

50 mg/mL; 10 mL Vial

ANDA # 40-333

Reviewer: Chandra S. Chaurasia

Gensia Sicor Pharmaceutical, Inc

Irvine, CA

Submission Date:

August 31, 1998

Review of a Waiver Request

BACKGROUND

- 1. The firm has requested a waiver of *in vivo* bioequivalence study requirements for its drug product, Fluorouracil Injection, USP, 50 mg/mL in 10 mL vial. The reference listed drug (RLD) is Fluorouracil Injection, 50 mg/mL in 10 mL vial manufactured by The Roche Laboratories, NDA #12-209.
- 2. The drug is indicated for the palliative management of carcinoma of the colon, rectum, breast, stomach and pancreas.

FORMULATION COMPARISON

Components and composition of the test and the reference products are as follows:

Comparison of Formulations						
Ingredient	Test Product	RLD				
	(mg/mL)	(mg/mL)				
Fluorouracil, USP	50	50 .				
Sodium hydroxide, NF	to adjust pH	to adjust pH				

COMMENTS

- 1. The drug is classified "AP" in the list of the "Approved Drug Products with Therapeutic Equivalence Evaluation".
- 2. The test drug product contains the same active and inactive ingredients in the same concentrations as the currently approved listed reference product and is intended solely for administration by injection.

3. The waiver of *in vivo* bioequivalence study requirements may be granted based on 21 CFR § 320.22(b)(1) of the Bioavailability/Bioequivalence Regulations.

RECOMMENDATION

The Division of Bioequivalence agrees that the information submitted by Gensia Sicor Pharmaceuticals, Inc. demonstrates that its Fluorouracil Injection, USP, 50 mg/mL in 10 mL vial falls under 21 CFR § 320.22(b)(1) of Bioavailability/Bioequivalence Regulations. The waiver of in vivo bioequivalence study for Fluorouracil Injection, USP 50 mg/mL in 10 mL vial of the test product is granted. From the bioequivalence point of view, the Division of Bioequivalence deems Gensia Sicor's Fluorouracil Injection, USP 50 mg/mL in 10 mL vial to be bioequivalent to the reference listed product, Roche's Fluorouracil 50 mg/mL, 10 mL vials.

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CMandra S. Chaurasia Division of Bioequivalence Review Branch I

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Concur	(/\$/	Date:	11/2/98
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